

Figure 1. The effect of the number of iterations on the accuracy of the proposed algorithm.

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CLASS A GROUP II						
A1AB_human	α_{1B} -adrenergic alpha 1B-AR	TMDI junction between TMDIII and IC2	63 FAIVGNILVIL A 142 CAISIDRYIGV A	IP / COS-7	(Scheer, Fanelli et al. 1997)	
A1AB_human	α_{1B} -adrenergic alpha 1B-AR	junction between TMDIII and IC2	143 CAISIDRYIGV K	IP / COS-7	(Scheer, Costa et al. 2000)	
A1AB_human	α_{1B} -adrenergic	TMIII	128 AVDVLGCTASI F	IP / COS-1	(Perez, Hwa et al. 1999)	
A1AB_human	α_{1B} -adrenergic	carboxyl end of IC3 TMV	293 REKKAÄKTIGI E 204 EERFYÄLPSSIG V	IP arachidonic acid release IP / COS-1	(Hwa, Garvin et al. 1997)	
A1AB_human	α_{1B} -adrenergic	C-terminal IC3	293 SREKKAÄKT X=19 different substitutions	PI / COS-7	(Kjelsberg, Cotecchia et al. 1992)	
A1AB_human	α_{1B} -adrenergic	C-terminus IC3	288 293 KFSREKKAÄKTIGI K H L	PI hydrolysis / rat fibroblast	(Allen, Leikowitz et al. 1991)	
A2AA_human	α_2 C10-adrenergic	C-terminal IC3 loop	373 (348?) EKRFTEVLAV X=F,A,C,E,K	adenylyl cyclase inhibition / HEK293	(Ren, Kurose et al. 1993)	
ACM1_human	alpha-2AAR muscarinic Hm1	C-terminal IC3 loop junction	360 SLVKEKKAARTLS A	PI / HEK(U293)	(Högger, Shogkley et al. 1995)	
ACM2-human	muscarinic acetylcholine M1 muscarinic acetylcholine M2	junction of IC3 and TMVI	390 KKVTRTTL1A 1-4 A inserted	IP production, inhibition of cAMP production / COS-7	(Liu, Blin et al. 1996)	

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CLASS A					
GROUP II					
ACM3_rat	m3 muscarinic (rat)	TMVI	507 TWTPYNIMVLVNT S	IP / COS-7	(Büml, Mutschler et al. 1994)
ACM5_human	muscarinic acetylcholine M3	N-terminus to TMII	chimera composed of m2 1-69 m5 77-445 m2 391-466	β -gal / NIH 3T3	(Burststein, Spalding et al. 1996)
ACM5_human	m5 muscarinic	TMVI	451 ALLA EITTW TPYNI MVLVST M L H C V S F T	β -gal; radioligand binding / NIH-3T3	(Spalding, Burststein et al. 1998)
ACM5_human	muscarinic acetylcholine M5	TMVI	451 ALLA EITTW TPYNI MVLVST M L H C V S F T	β -gal; radioligand binding / NIH-3T3	(Spalding, Burststein et al. 1997)
ACM5_human	m5 muscarinic	junction of TMVI and EC3	465 YNIMVLVSTFCDCV X=V,F,R,K,+more	β -gal; radioligand binding / NIH-3T3	(Spalding, Burststein et al. 1997)
BIAR_human	β_1 -adrenergic	C-terminus	389 RKAFOGLCCA R	adenylyl cyclase; agonist binding / CHW	(Mason, Moore et al. 1999)
B2AR_human	β_2 -adrenergic	C-terminal IC3 loop	266 272 FCLKEHKALKTIGI SR K A	adenylyl cyclase activation; agonist binding affinity / COS-7 or CHO	(Samama, Cotecchia et al. 1993); (Lefkowitz, Cotecchia et al. 1993)
DADR_human	dopamine D1A	carboxyl terminal IC3	264 SFKMSFKRETQVLT I K 288 from D1B receptor APDTSIKKETKVLKT	adenylyl cyclase; cAMP accumulation / HEK293	(Charpentier, Jarvie et al. 1996)
DADR_human	dopamine D1	TMVI	286 FVCCWLPFFIL A	CAMP accumulation / COS-7	(Cho, Taylor et al. 1996)
HH2R_rat	histamine H ₂	IC2	115 FMISLDRYCAV N,A	CAMP production / HEK-293	(Alewijse, Timmerman et al. 2000)

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File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS A GROUP III					
OPSD_human	opsin rhodopsin	TMII TMIII	90 FMVLGGFTSTLY D 113 GCNLEGGFPAT Q 292 296 MTIPAPFAKSAIY E G, E, M 297 Ala neutral a.a converted to carboxylate and competes with ¹¹³ Glu for salt bridge with ²⁹⁶ Lys	transducin; phosphorylation by rhodopsin kinase / COS	(Rim and Oprian 1995)
OPSD_human	opsin rhodopsin	TMIII	134 VLAIERVVV I, Q, S	transducin; radioligand binding / COS	(Acharya and Karnik 1996)
OPSD_human	opsin rhodopsin	TM6	257 RMVILMIVIAFL Y, N	transducin, GTPγS uptake / COS	(Han, Smith et al. 1998)
OPSD_human	opsin rhodopsin	plus TM3 TMVII	plus G113Q 296 PAFFAKSAIY G X=E,M natural mutants + 10 different a.a. substitutions disrupts critical salt bridge between ²⁹⁶ Lys(TMVII) and ¹¹³ Glu(TMIII)	transducin; radioligand binding / COS	(Govardhan and Oprian 1994); (Cohen, Yang et al. 1993)
		IC2	134 VLAIERVVV Q		(Cohen, Yang et al. 1993)

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File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS A GROUP V					
AG2R_rat	AT _{1A}	TMIII	111 ASVSFNLVASV A disrupts human (TMIII) - 292TYR (TMVII) interaction	phospholipase C, IP production / COS-7	(Grobowski, Maigret et al. 1997)
AG2R_rat	Type-1A angiotensin II	C-terminus of TM7	305 LFYGFYGGKPK Q	IP production / HEK-293; intracellular Ca ²⁺ mobilization / CHO	(Parnot, Bardin et al. 2000)
FM1R_human	Type-1A angiotensin II formylmethionyleucylphenylalanine (FM1PR)	other multiple mutations IC1	51 LVYVAGFRMTHTYTTISYLNKAVA LVVWVTAFEAKRKTINAIWFLNLAVA (K above conflicts with SWISS-PROT database)	PI production; phospholipase C stimulation / COS-7	(Amatruda, Draga-Graonic et al. 1995)
IL8B_human	interleukin-8 receptor B	IC2	138 ACISVDRLAIVH V	IP production; Ca ²⁺ mobilization and actin polymerization / NIH 3T3	(Burger, Burger et al. 1999)
LSHR_human	CXCR-2 chemokine	IC3	564 MATNKDTKIACK G	CAMP production / HEK293	(Kudo, Osuga et al. 1996)
LSHR_human	luteinizing hormone (LH)	IC3	578 ILIFTDFTCMA G	CAMP production / COS-7	(Shenker, Laue et al. 1993)
LSHR_human	luteinizing hormone (LH)	TMVI	571 577 KIAKKMAILIFTDFTCM I I	CAMP production / COS-7	(Kosugi, Van Doo et al. 1995)
LSHR_human	luteinizing hormone (LH)	TM6	556 ILIFTDFTCMA G, Y	CAMP production / HEK 293T	(Bradbury, Kawate et al. 1997; Bradbury and Menon 1999)
OPRD_mouse	delta opioid receptor	TM3	128 KVLSTIDYYNMF A, K, H	adenylyl cyclase inhibition / COS-7	(Cavalli, Babey et al. 1999)
OXVR_human	oxytocin	IC2	137 LMSLDRCIAIC A	IP production / COS-7	(Franeli, Barbier et al. 1999)

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PAFR_human	platelet-activating factor (PAF)	C-terminus of IC3	231 EVKRALIMVCTVLAV R	IP production / COS-7	(Parent, Le G ull et al. 1996)
PAFR_human	platelet-activating factor (PAF)	TMIII	100 CLFIIINITYCSV A	arachidonate release, IP production, adenylyl cyclase inhibition / CHO	(Ishii, Izumi et al. 1997)
PE23_human	prostaglandin E ₂ , EP3III EP3IV	C-terminal tail	360 FCOEFEFGN FCOMRRRLRQBEFGN ↑truncated	inhibition of adenylyl cyclase / CHO-K1	(Jin, Mao et al. 1997)
PE23_mouse	prostaglandin E ₂ , EP3	carboxyl-terminal tail	336 KILRKFCQIRDHT (3α) MNNHL (3β) ↑truncated	inhibition of adenylyl cyclase / CHO, stably expressed	(Hasegawa, Negishi et al. 1996)
THRR_human	thrombin	EC2 loop	259 CHDVLENTLEGGYAYY DLKD KDF I 486 YNNHAIDWQTG F, M 568 YAKVSI ^T CLPMD	⁴⁵ Ca ²⁺ efflux, PI hydrolysis, reporter gene induction / COS-7	(Nanevicz, Wang et al. 1996)
TSHR_human	thyrotropin (TSHR)	EC1	486 YNNHAIDWQTG F, M 568 YAKVSI ^T CLPMD	inositol phosphate--diacylglycerol cascade / COS-7	(Parma, Van Sande et al. 1995)
TSHR_human	thyrotropin (TSHR)	TMIII	509 ASELSYTLTV A	adenylyl cyclase activation / COS-7	(Duprez, Parma et al. 1994)
	thyroid stimulating hormone	TMVII	672 YPLNSCANPFL Y		
TSHR_human	thyrotropin (TSHR)	TMV	597 VAFVIYCCCHV L	cAMP formation / COS-7 cells	(Esapa, Duprez et al. 1999)
TSHR_human	thyrotropin (TSHR)	TMVII	677 CANPFLYALFT V	cAMP formation / CHO cells	(Russo, Wong et al. 1999)
TSHR_human	thyroid stimulating hormone	IC3	613 VRNPOYNPGDKDTKIAK deletion	cAMP formation / COS-7	(Wonerow, Schoneberg et al. 1998)

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Sequence

TSHR_human	thyrotropin (TSHR)	IC3/TMVI	623 KPTKIAKPMAVLIPTDFICM V 136 LAFTLDKRRAL A	632 I	cAMP activation / COS-7	(Pasciute, Tonacchera et al. 1994)
V2R_human	thyroid stimulating hormone vasopressin V2	IC2			cAMP formation / COS-7	(Morin, Cotte et al. 1998)

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[illegible][illegible]

File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS D 074283 RCB2	pheromone	TM6	229 PLSAVQIVLGT P	heterologous yeast assay	(Olesnick, Brown et al. 1999)
<i>C. cinereus</i> STE2_yeast	pheromone α -factor	TM6	258 QSLVPSIIFI LL	<i>lacZ</i> reporter gene	(Konopka, Margani et al. 1996)
STE2_yeast	pheromone α -factor	double mutations TM5 and TM6	223 MSFVLYVKKILAIR C C 247 251 DSFHILLIKCOSIL CC CC	<i>lacZ</i> reporter gene / yeast	(Dube, DeCostanzo et al. 2000)
STE3_yeast	pheromone α -factor	IC3	double mutations 194 DVRDILHCTNS	β -galactosidase	(Boone, Davis et al. 1993)
STE2_yeast	pheromone α -factor	TM6	253 258 LIMSCQSLVPSIIFI L LP	β -galactosidase	(Sommers, Martin et al. 2000)

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A Point Mutation Enhances MC-4 Receptor Constitutive Activity

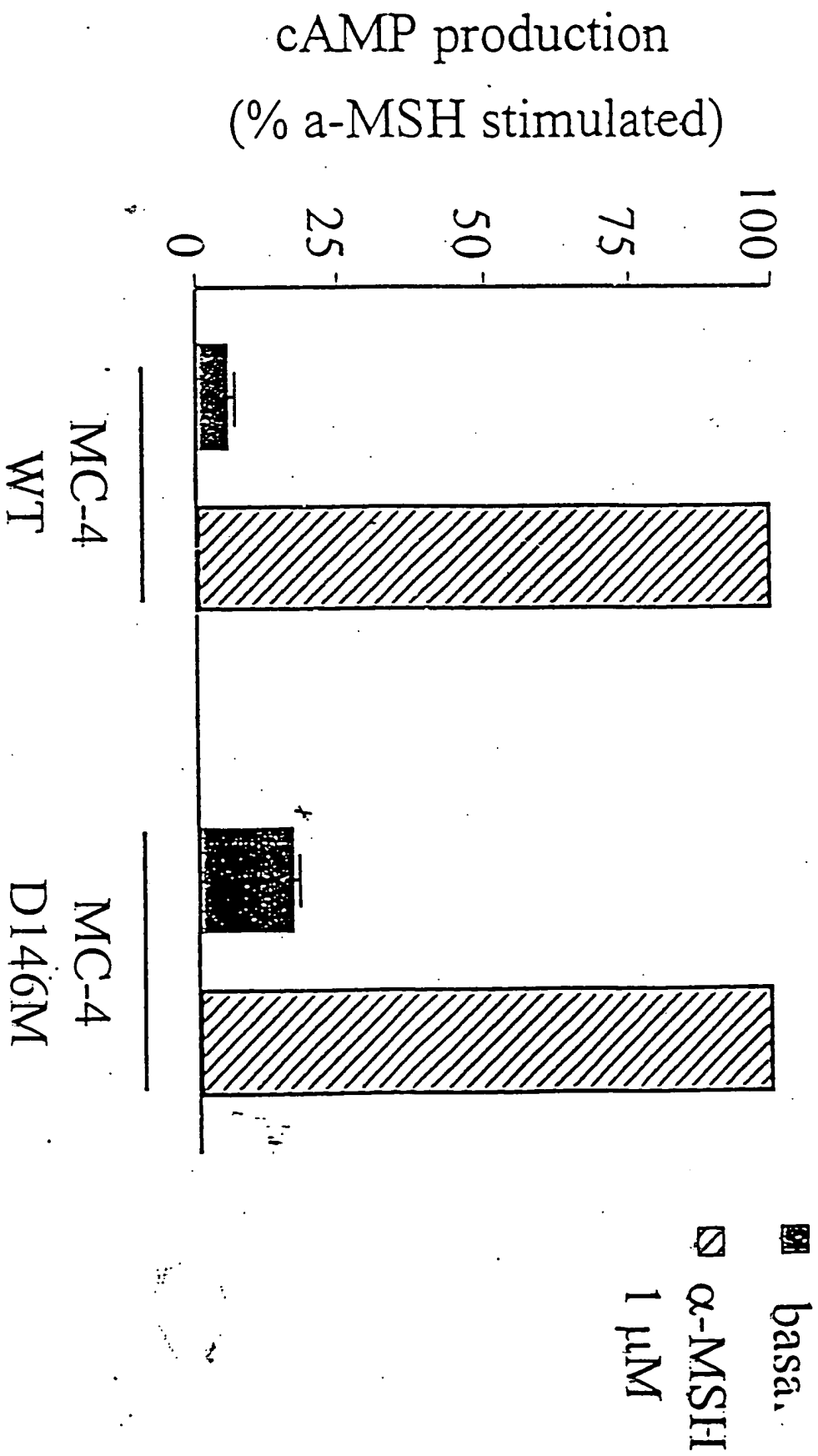


Figure 2

Light Emission Induced by the WT CCK-BR vs. a Constitutively Active Mutant

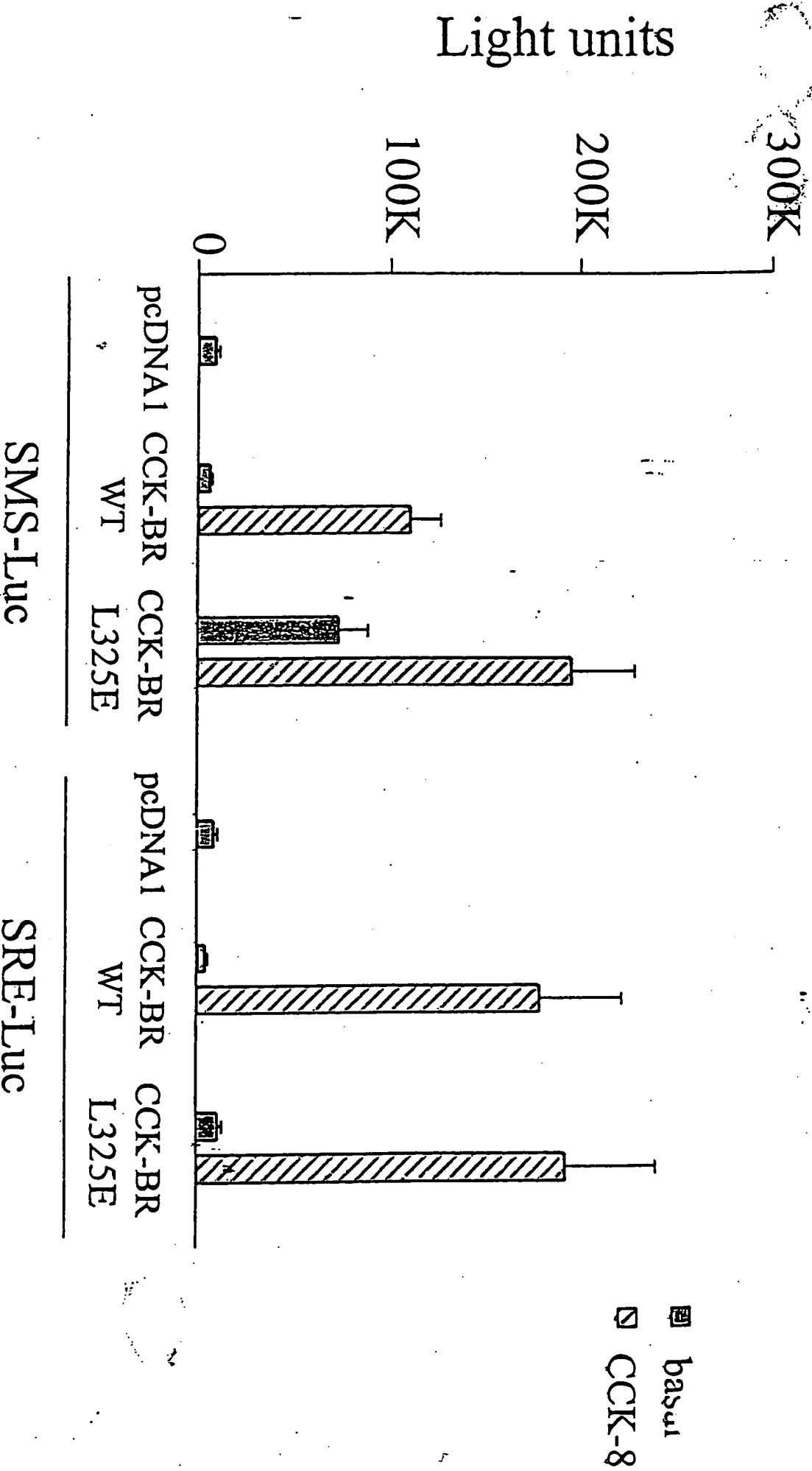


Figure 3

A Point Mutation Confers Constitutive Activity to the Rat μ Opioid Receptor

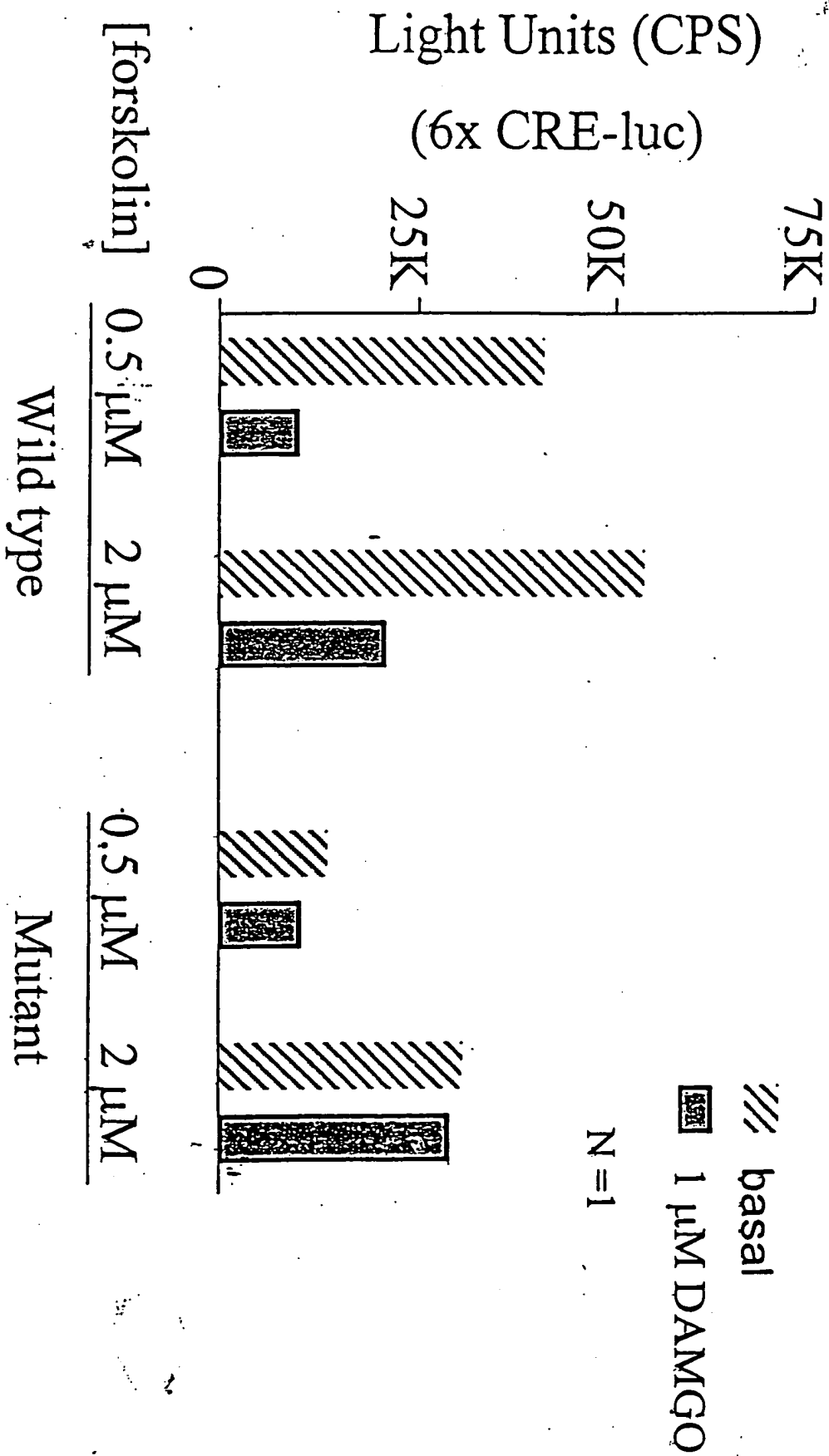


Figure 4

Forskolin Stimulated HEK293 Cells Transfected With pcDNA1 and a CRE-luc Construct

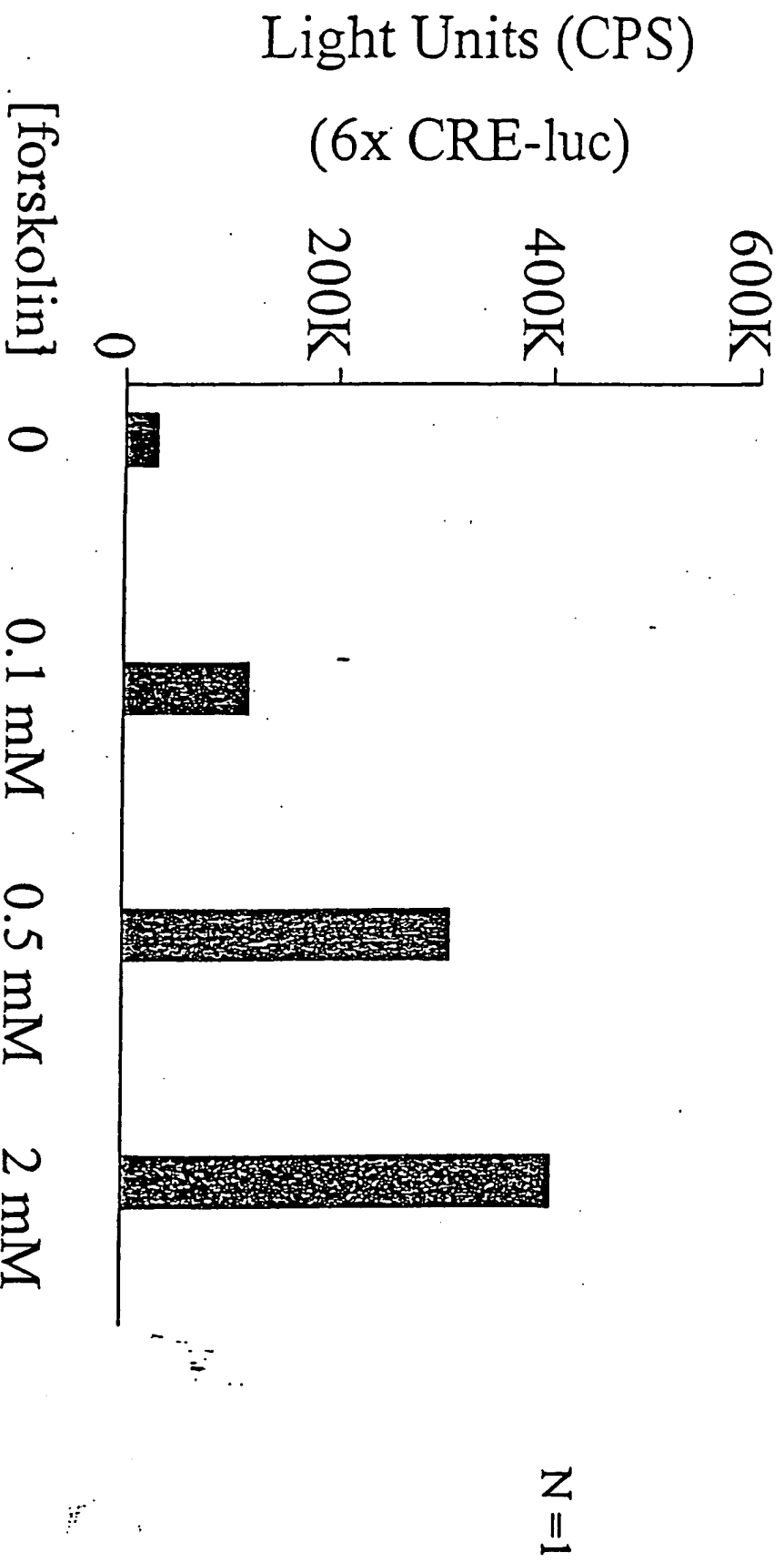


Figure 5

The Rat μ Opioid Receptor Signals Through $G_{\alpha i}$

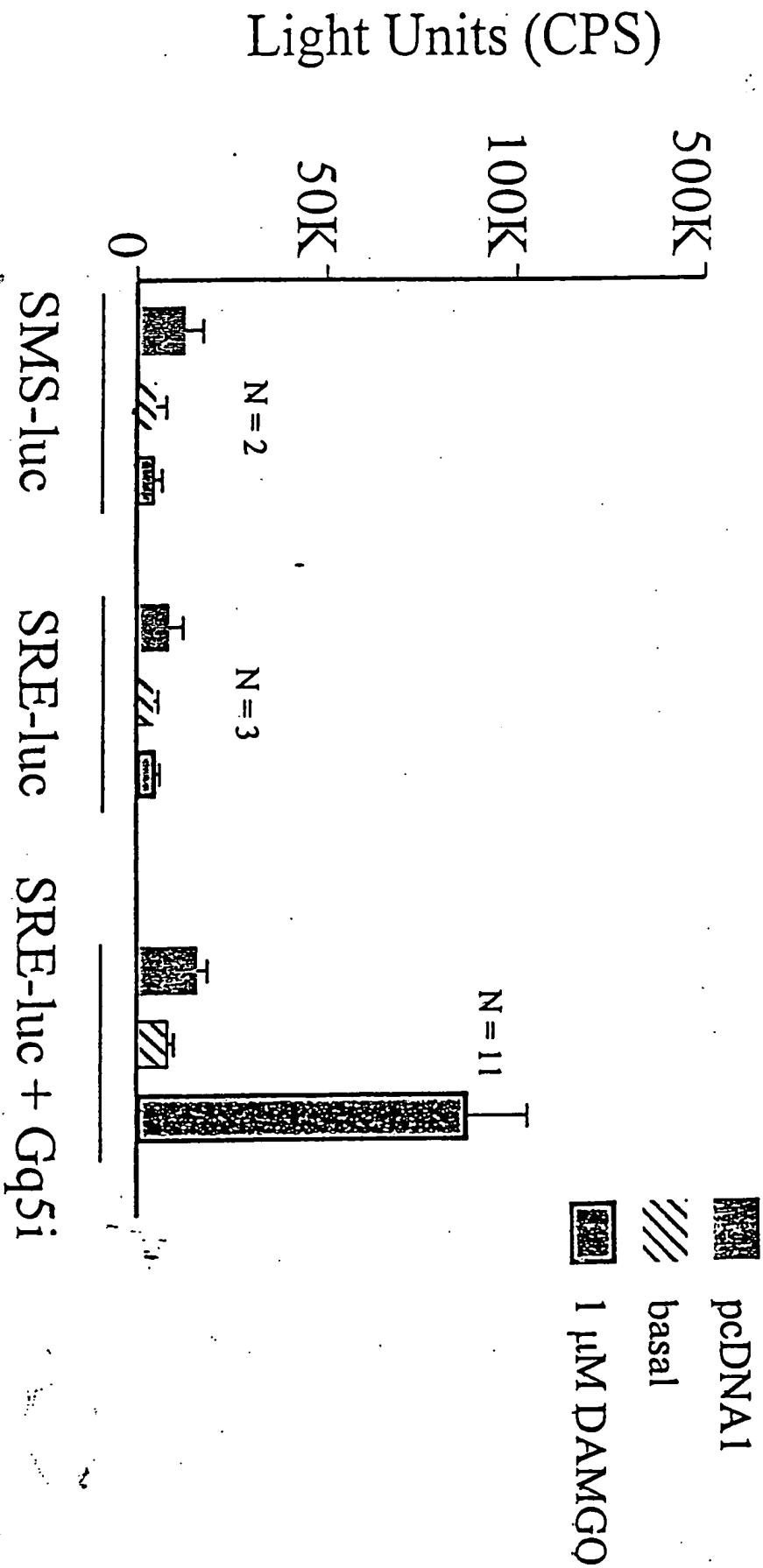


Figure 6

A Point Mutation Confers Constitutive Activity to the Rat μ Opioid Receptor

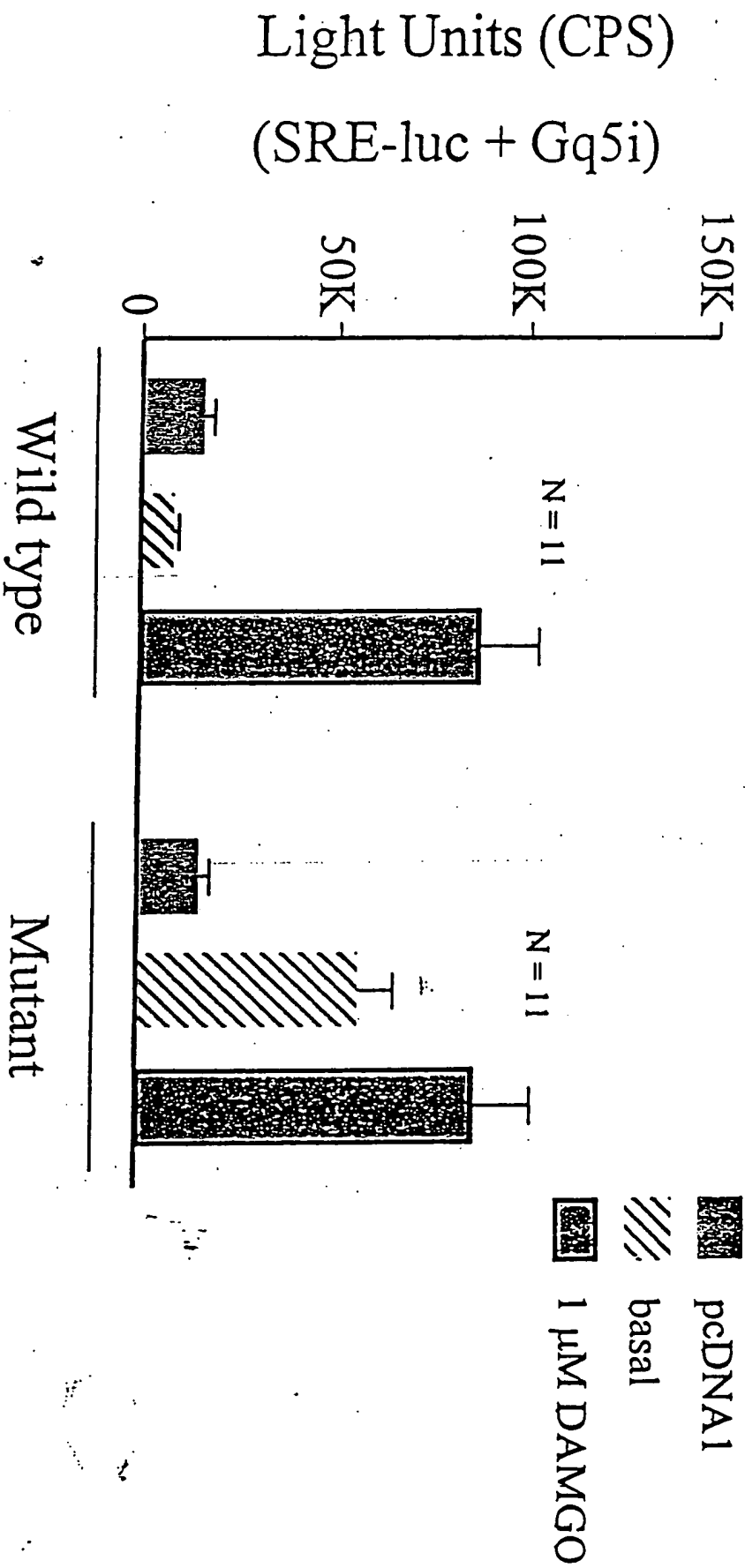


Figure 7

Target Residues Within Class I GPCRs

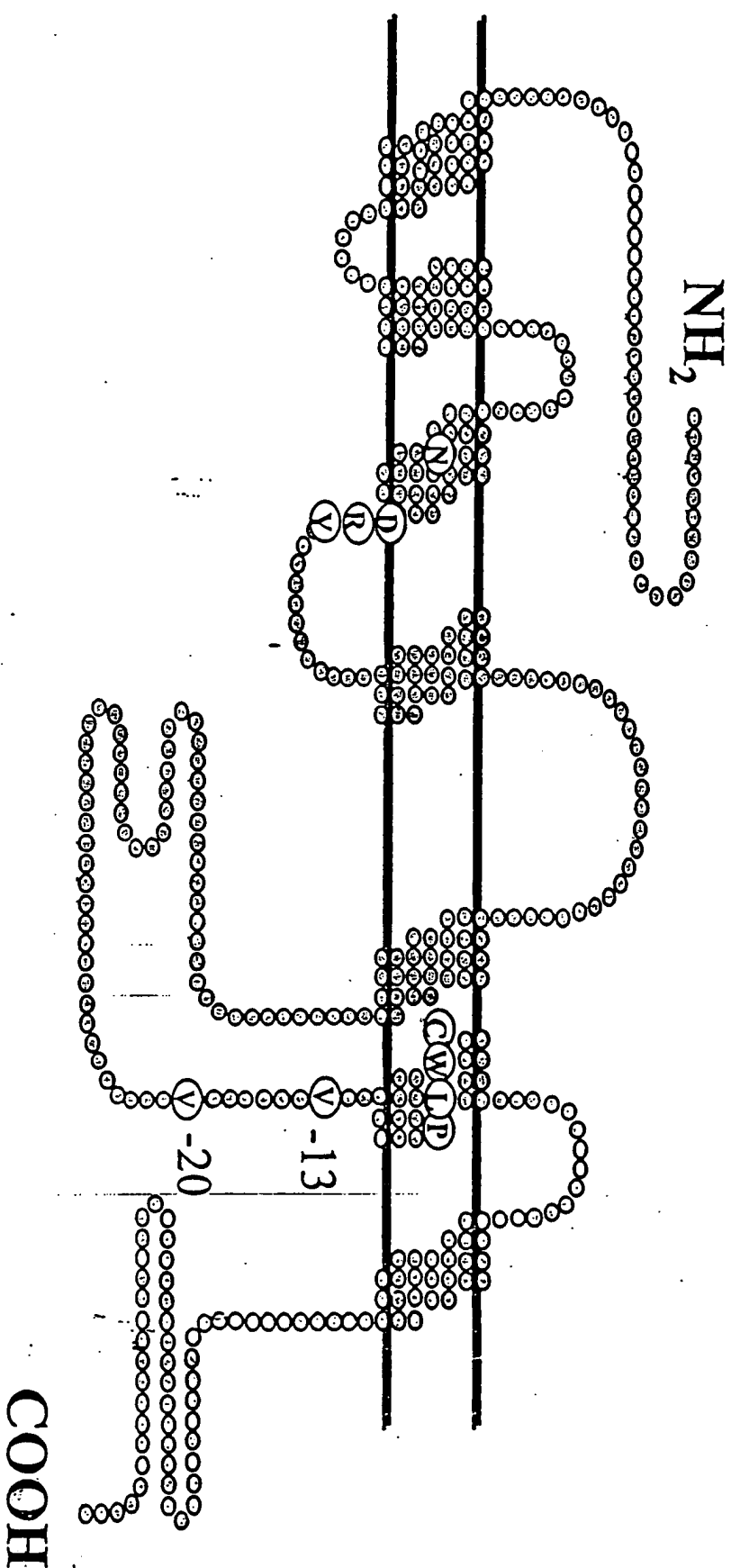


Figure 8

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TMD III Asn (-14 from DRY) is a Target for Mutation Induced Constitutive Activity

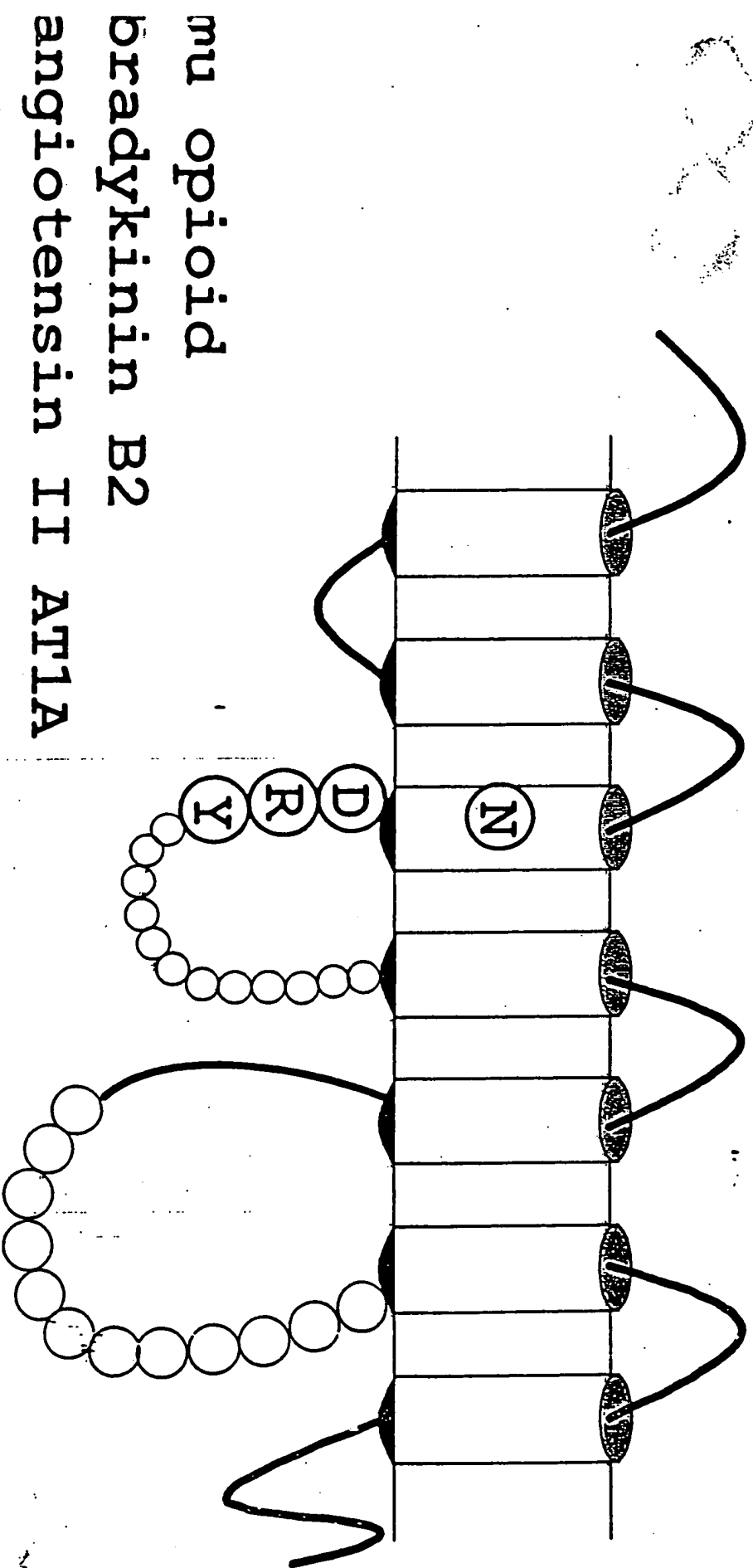


Figure 9

The 'DRY' Motif is a Target for Mutation Induced Constitutive Activity

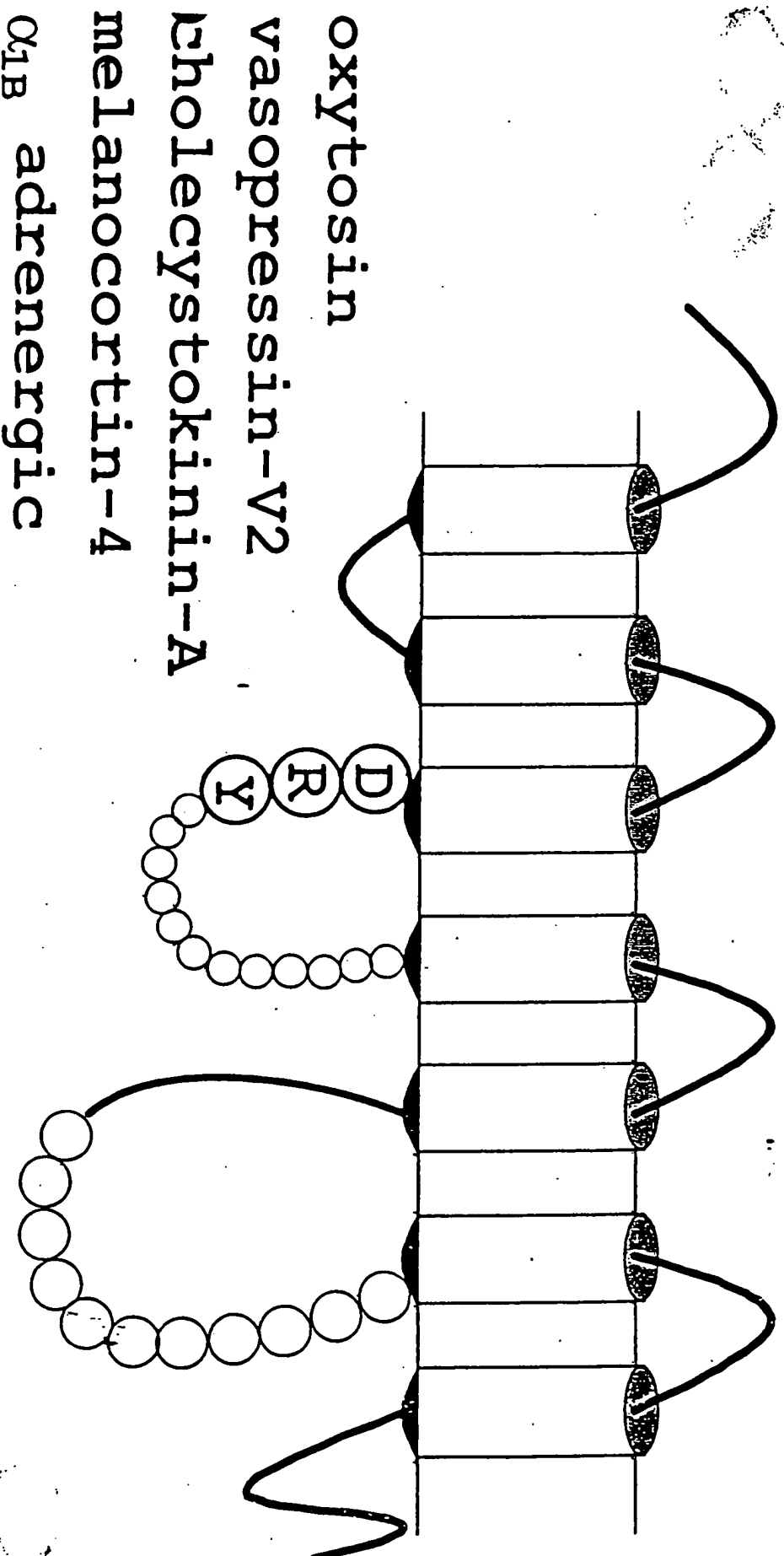


Figure 10

A Point Mutation Enhances MC-4 Receptor Constitutive Activity

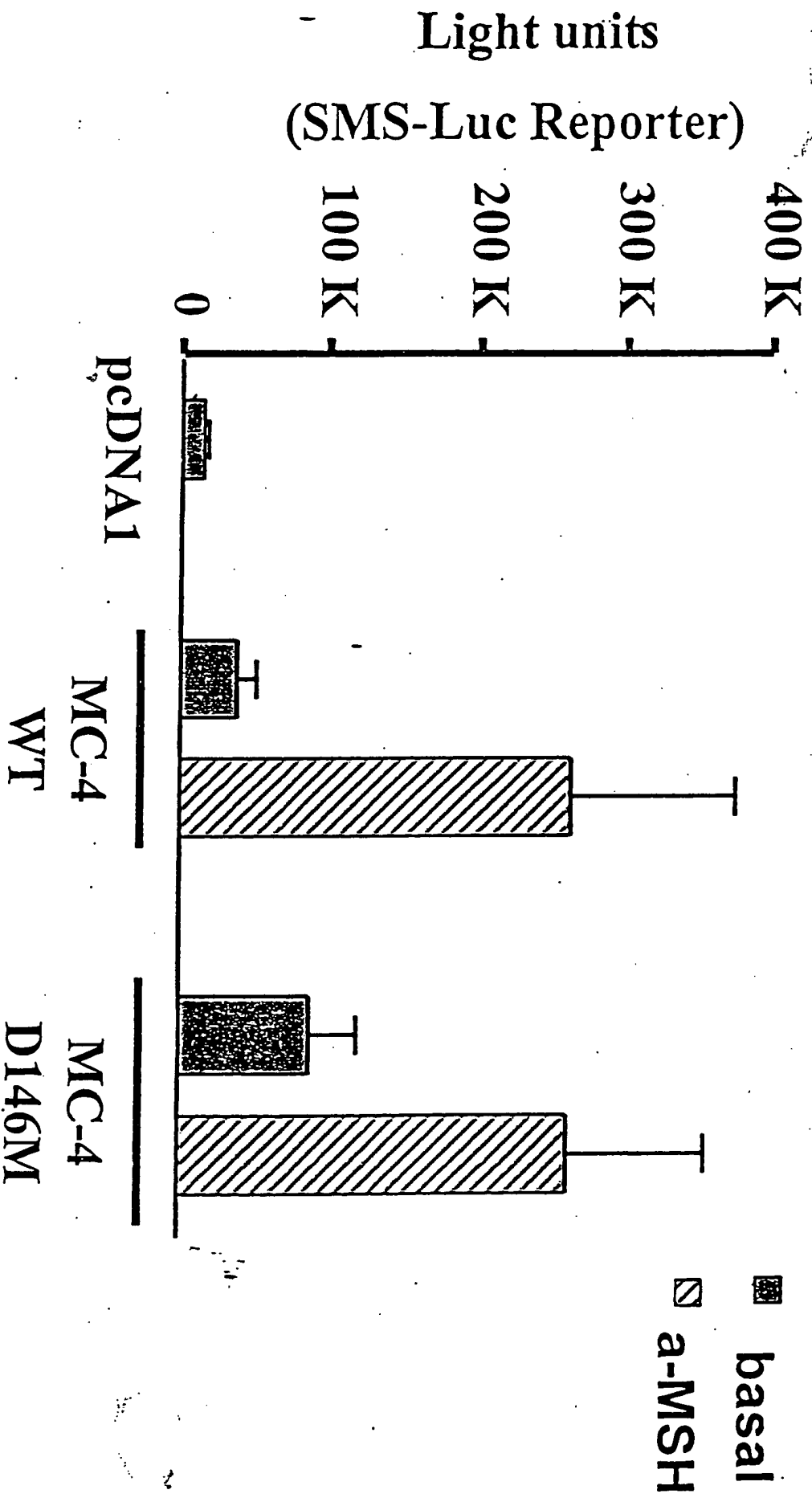
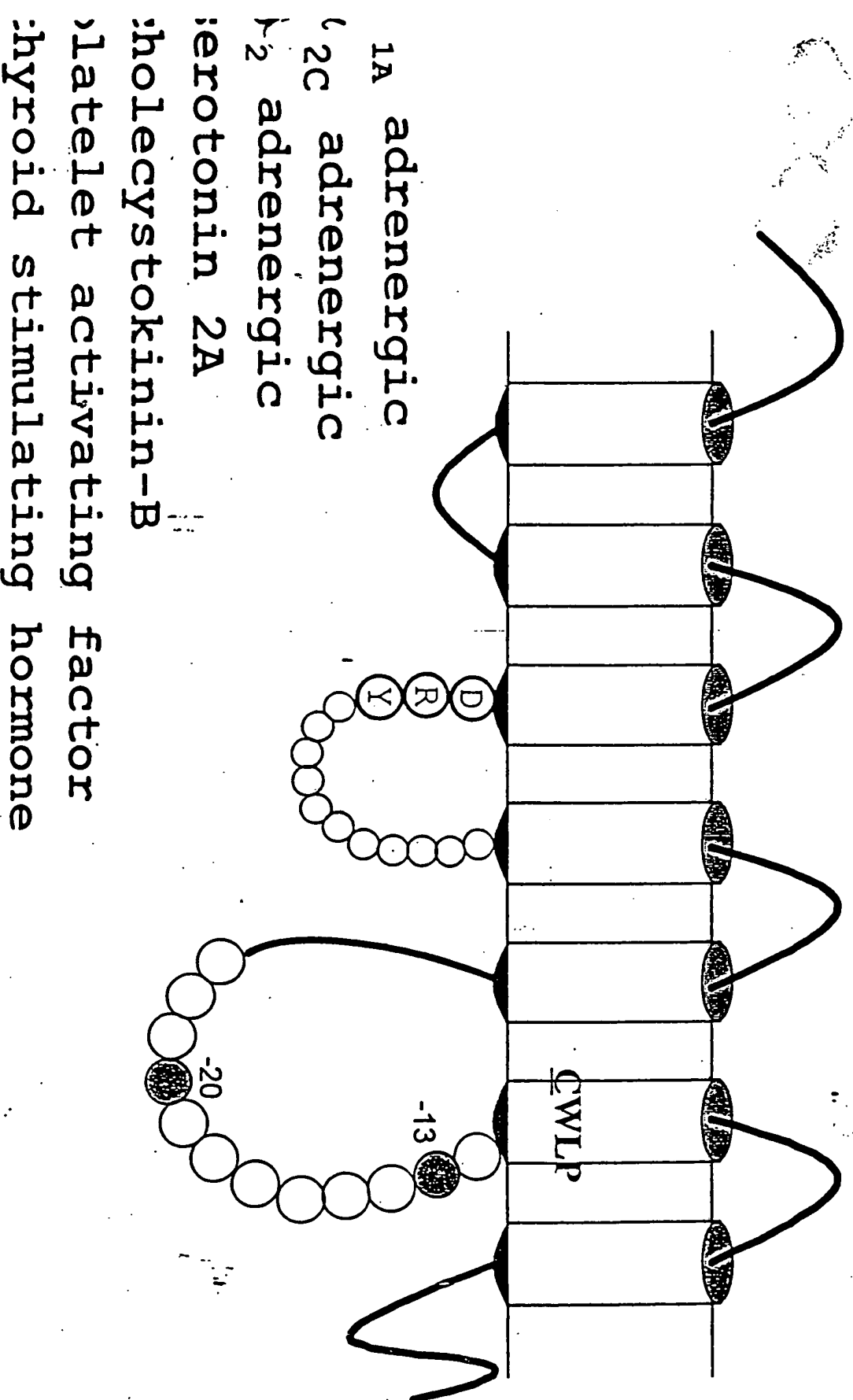


Figure 11

The -13 Position is a Target for Mutation Induced Constitutive Activity



[illegible]

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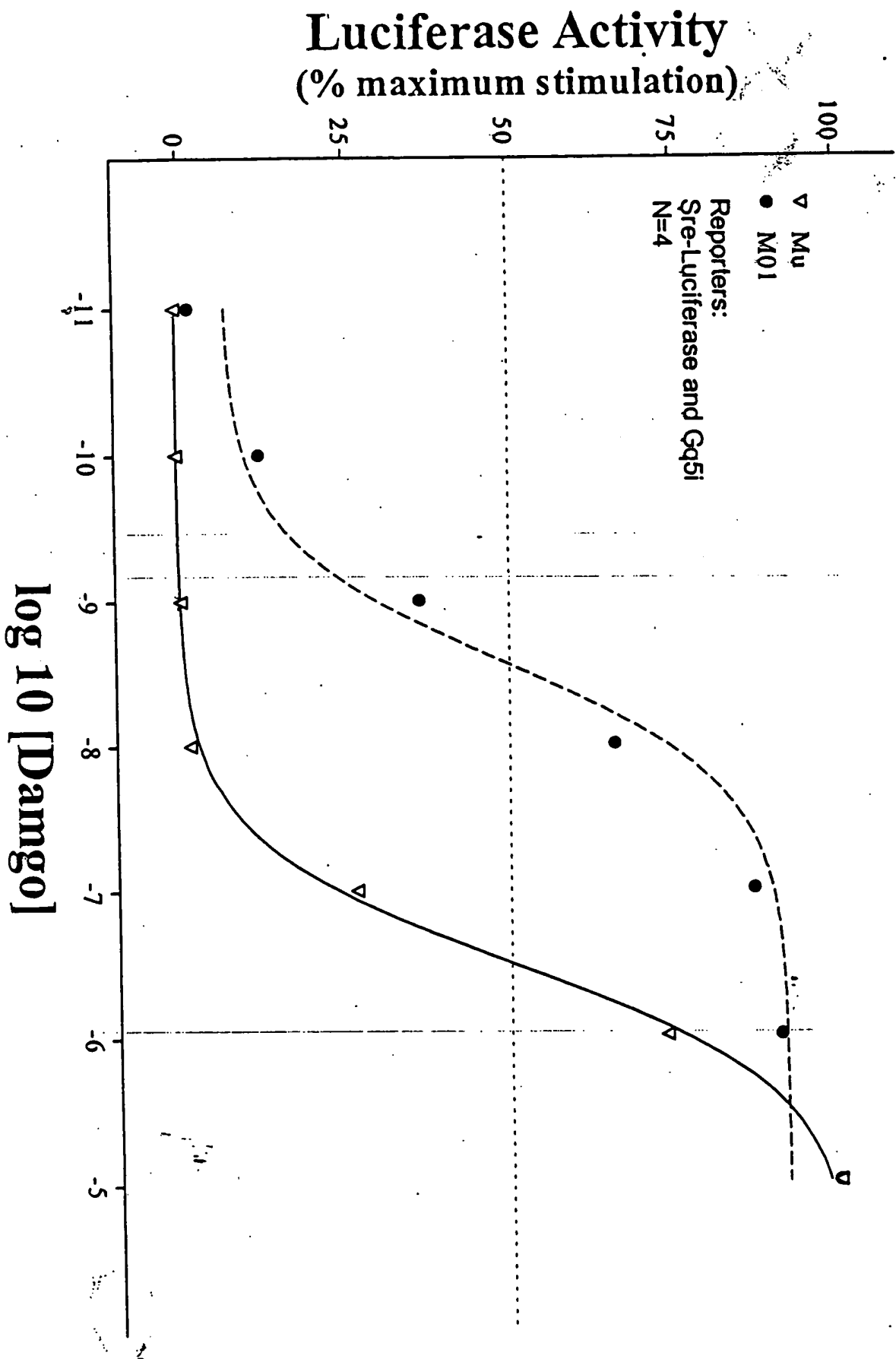


Figure 15

An Intracellular Point Mutation Results in Loss of Ligand-Induced Function

IP Production / ³H Inositol incorporation

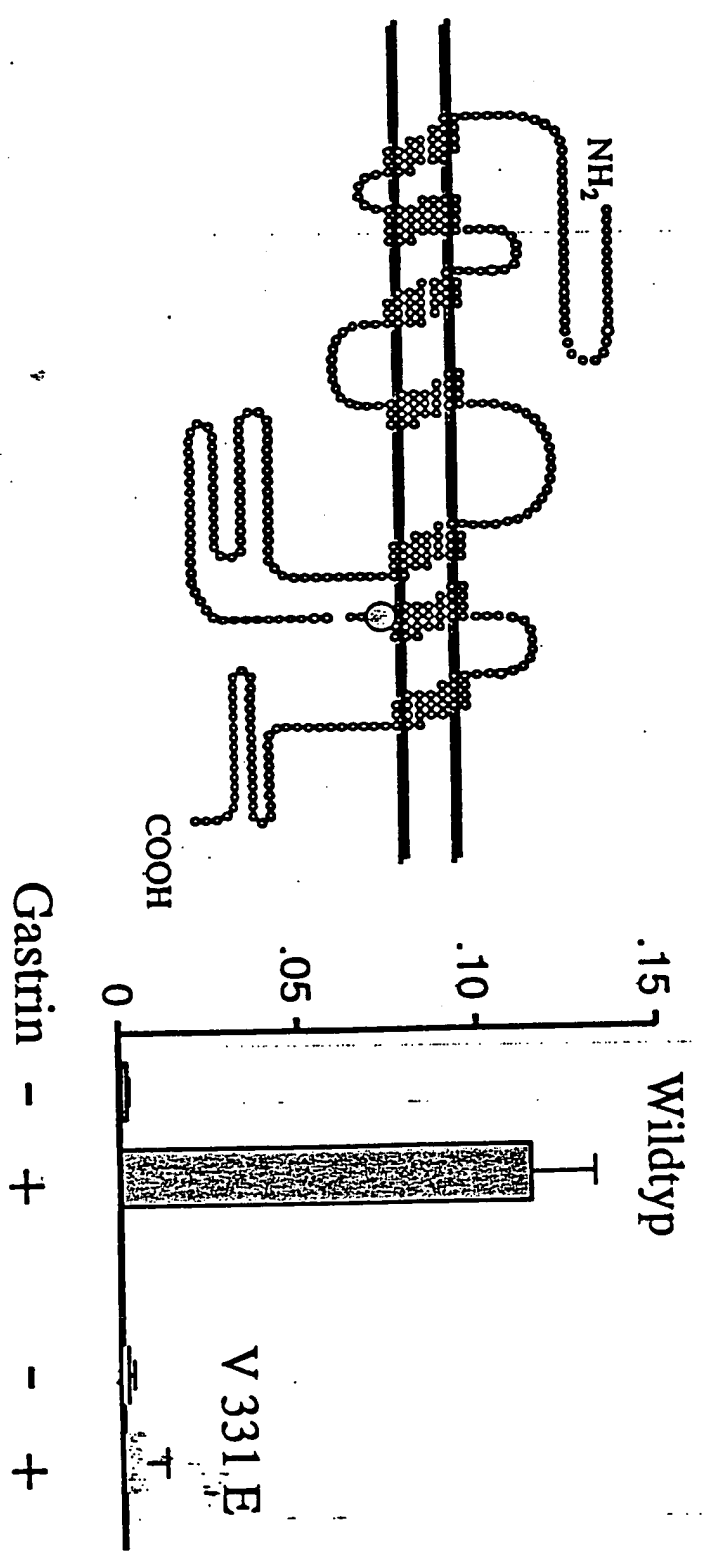


Figure 16

